

# Impact of Natural Selection on Lignin and Cellulose Candidate Genes in a Natural Population of *Eucalyptus urophylla*

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Received 3 August 2014; revised 6 September 2014; accepted 21 September 2014

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## Abstract

Wood plays a major role in land ecosystems and in human activity. Better understanding the genetic basis and evolutionary implication of wood variability are thus key issues with both ecological and economical implications. The present paper addresses the question of the extending and the nature of natural selection on wood related genes in *Eucalyptus urophylla*, a tropical tree species with key economical importance. We conducted a genetic study on an *E. urophylla* population from Timor Island using a set of 17 SSR characterized on a main sample of 43 individuals and six candidate genes sequenced on a subset of 18 individuals. The candidate genes include three cellulose synthase genes (*EuCesA1*, *EuCesA2* and *EuCesA3*), and three genes involved in lignin synthesis (*EuCAD2*, *EuC4H1* and *EuC4H2*). Based on SSR data, the investigated population appeared to have no structure and have undergone past population expansion. Accounting for this demographic history, we were able to draw neutral expectation for polymorphism distribution on candidate genes and to determine their potential selective status. We hence identified two gene portions exhibiting unexpected polymorphism pattern, consistent with natural selection imprint.

## Keywords

*Eucalyptus urophylla*, Adaptation, Demography, Approximate Bayesian Computation, Candidate Genes

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## 1. Introduction

Wood is a structural tissue resulting from the secondary growth of vascular cambium in tree stems and roots. It is a natural composite of cellulose fibers embedded in a lignin matrix which provides trees with physical support, water and nutriment transport between roots and leaves and nutriment storage. Wood is quantitatively and qualitatively a key component of land ecosystems and its properties further determine biomass decomposition rates, playing a key role in carbon and nitrogen cycle [1] [2]. It is also a very important resource for humans, being used to produce paper, charcoal, furniture and many other facilities. Wood first evidence dates back from early Devonian, 407 million years ago [3] and, since then, its chemical and physical properties have highly evolved. This variation exhibits complex patterns, which globally appear to be the results of functional constraints, phylogenetic conservatism and environmental adaptation [4]. Focusing at intra-specific level, wood traits exhibit important variability with high heritability [5], raising the question of their particular role in species adaptation and the selective forces acting on them. Wood is a complex structure with thousands of genes involved [6]-[12] but an increasing number of studies have managed to pinpoint gene SNPs associated to wood property variability in natural populations of several tree genus, including *Eucalyptus* and allied species [13]-[17]. On the other hand, the question of the natural selection pressures acting on wood related genes has been rarely addressed, in particular at intra-specific level. Wood related genes are generally considered to be under purifying or balancing selection, but only few examples have been reported so far [18] [19].

The present paper attends to decipher the selective forces acting on six full length wood synthesis candidate genes in *Eucalyptus urophylla* S T Blake. *E. urophylla* is the only *Eucalyptus* species not naturally found in Australia and has a relatively restricted area, the Lesser Sunda Islands where it occurs as disjunct populations across seven islands, with main stands in Timor and Wetar [20]. *E. urophylla* is one of the main *Eucalyptus* species used for industrial plantations in tropical regions and it is also described to be one of the most morphologically variable species among *Eucalyptus* [21], being thus a very good model to study adaptation. *E. urophylla* grows from sea level up to 3000 m and displays dramatic phenotypic differences as for growth and shape as well as for fruit size and bark morphology [20] [22].

We focus here on a set of lignin (*EuC4H1*, *EuC4H2* and *EuCAD2*) and cellulose synthesis (*EuCesA1*, *EuCesA2* and *EuCesA3*) gene set for which association with wood related traits has been already investigated in *E. urophylla* [17]. *Cellulose synthesis (CesA)* genes encode components of an enzyme complex embedded in the cellular membrane [23] and characterized by a “rosette” like structure [24]. This enzyme complex catalyzes the last biochemical reaction of the cellulose synthesis and synthesizes six  $\beta$ -1,4-glucan chains that cocrystallize to form a 36-chain microfibril [25]. While *CesA* genes can be expressed either in primary tissues or in secondary tissues [26], *EuCesA1*, *EuCesA2* and *EuCesA3* were chosen because in *E. globulus* Labill., a species closely related to *E. urophylla*, the three orthologous *CesA* genes are specifically and strongly expressed in wood [27]. *C4H* and *CAD2* are two enzymes that play an important role at the beginning and the end of the lignin biosynthesis pathway. *Cinnamate 4-hydroxylase (C4H)* belongs to the core reactions of phenylpropanoid metabolism. It catalyzes the hydroxylation of cinnamic acid to 4-hydroxycinnamate (or p-Coumaric acid), which is the second step of the phenylpropanoid pathway [28]. Evidences suggest that *C4H* has been duplicated before monocot and dicot separation and can be found in a single or several copies in plants [28]-[30]. Finally the *CAD2* enzyme belongs to the specific lignin pathway. The *Cinnamoyl CoA Reductase (CCR)* converts hydroxycinnamoyl CoA esters to their corresponding aldehydes, which are converted in the monomeric precursors of lignin by the *Cinnamyl Alcohol Dehydrogenase (CAD2)* [28].

In order to characterize the selective forces acting on *E. urophylla* wood gene at intra-specific level, we sequenced the six candidate genes on 18 individuals from Timor Island and investigated the imprint of natural selection on their polymorphism pattern. Since demographic events can generate polymorphism pattern similar to what is expected under selection at the whole genome scale, it is important to account for it in order to avoid false positives. However, the demographic history of *E. urophylla* is not well known. Previous studies showed no [22] or weak population structure [31] at species range level. Evidence for bottleneck was specifically investigated and was found on chloroplast markers [32] but not on nuclear SSR data. To clarify demographic history of our *E. urophylla* sampled region, we used a set of 43 individuals genotyped for 17 SSR markers to evaluate the fit of several simple demographic scenarios using Approximate Bayesian Computation (ABC) framework. ABC is a flexible class of Monte-Carlo algorithms used to perform model-based inference [33] and relying on summary statistics of the data, making it much less computationally demanding than full likelihood relying ap-

proaches. The demographic history being elucidated on SSR data, the results served as a basis to build an ad-hoc expected distribution for sequence polymorphism pattern accounting for demographic history under selective neutrality. We used this ad-hoc distribution along with other neutrality test to evaluate selective status of the candidates.

## 2. Material and Methods

### 2.1. Sampling and DNA Extraction

To perform our study, we have selected samples from the East Timor (**Figure 1** and **Table 1**). No endangered or protected species were involved in this study. Seeds of *Eucalyptus urophylla* were collected between in 1973 and preserved in the CIRAD Genetic Forestry Laboratory. At the time of the sampling, East Timor was a Portuguese territory and sampling authorization was obtained from the military governor (SUPDOC). As GPS technology was not available at the time of the sampling, coordinates were estimated using a 1/50,000 topographic map. The seeds had been stored in a cold room at 4°C and 30% humidity. In 2003, some seeds from this collection were planted. The plantlets were maintained in humid tropical nursery at the following conditions: 28°C day temperature, 25°C night temperature, 70% humidity, pH4 substrate. Leaves were collected on 4 months old plantlets, dried in silica gel and stored at room temperature until DNA extraction. To perform our study, we have selected samples from the East of Timor Island (**Figure 1** and **Table 1**). We concentrate on Timor Island because the *E. urophylla* stands growing there are bigger and likely more stable over time than in other islands [20]. DNA was extracted from 43 individuals according to a modified protocol adapted from Gawel & Jarret [34] and Saghai-Marooof [35]. Each individual is coming from a different mother tree. All the 43 genotypes were used for the microsatellite analysis while 18 individuals were used for gene sequencing and gene polymorphism studies.

### 2.2. Microsatellite Genotyping

A total of 17 microsatellite loci were used for this study. They were all previously described: EMB18 [36]; EMB22, EMB27, EMB30, EMB32, EMB33, EMB37, EMB38, EMB42, EMB44, EMB47, EMB52, EMB56, EMB63 [37]; FMRA1, FMRA4, FMRA5 [38]. They were selected according to several criteria: specificity of the amplification, high polymorphism level, belonging to different linkage groups. The genotyping was per-



**Figure 1.** Sample location map. *E. urophylla* provenances sampled in this study are materialized on the map by pins associated with their number, as given in **Table 1**. Detailed geographical information about provenances is presented in **Table 1**. These geographic coordinates were obtained using a 1/50,000 topographic map. Map is similar to Open Street Map (<http://www.openstreetmap.org>) but not identical and is therefore for representative use only.

**Table 1.** Geographical data of the sampled individuals.

Num	Microsatellites	Gene	Location	Lat, Long	Alt
1	6	3	Pila-Paria, North Ermera	8°39'S, 125°28'E	488
2	7	1	Betularam, Mount Baudoe, Remexio	8°35'S, 125°45'E	517
3	4	0	AI-Betoulun, East Remexio	8°37'S, 125°41'E	902
4	7	0	Maulau, North West Turiscai	8°46'S, 125°40'E	950
5	4	0	Mano Mera Lolo	8°46'S, 125°34'E	1030
6	4	2	Lebo-Meta, Mount Berelico, Remexio	8°38'S, 125°44'E	1109
7	7	3	Rairema	8°48'S, 125°33'E	1146
8	10	0	Slaur-Lala, South West Remexio	8°40'S, 125°39'E	1147
9	9	3	Fatuc Hun, South West Remexio	8°38'S, 125°37'E	1193
10	9	0	Foho-Hua, Mount Aibali, Turiscai	8°49'S, 125°42'E	1269
11	6	1	Al Fero, South Maubisse	8°54'S, 125°36'E	1344
12	7	5	Flecha, South Maubisse	8°53'S, 125°36'E	1778

Num: location number reported on [Figure 1](#) map; Microsatellites: number of individuals used for the microsatellite analysis; Gene: number of individuals used for the gene sequence analysis; Lat, Long: latitude, longitude; Alt: altitude (expressed in meters). Latitude and longitude were estimated with a 1/50,000 topographic map.

formed on the 43 individual set and 3 DNA controls with known allele size were added to each 96 well plates. The PCR amplification was performed by multiplexing 2 primers. Five microliters of QIAGEN multiplex mix, 0.08  $\mu$ M of both forward primer with 5'-tail-end M13 (CACGACGTTGTAAAACGAC), 0.10  $\mu$ M of both reverse primer, 0.10  $\mu$ M IRDye fluorescent-labelled M13-primer (700 or 800 nm) and 5.0 ng of genomic template DNA. A touchdown cycling programme was used: 95°C for 15 min, 67°C for 1.5 min, 72°C for 1 min, followed by eight cycles of 94°C for 30 s, 65°C for 1.5 min with 2°C decrease at each cycle, 72°C for 1 min then 24 cycles at 94°C for 30 s, 51°C for 1.5 min, 72°C for 1 min, and a final extension of 60°C for 30 min. Amplified fragments were analysed at 700 and 800 nm by electrophoresis on an IR2-DNA analyzer (LI-COR 4200 sequencer) at the Montpellier Languedoc-Roussillon Genopole genotyping platform. Allele scoring was done with SAGA software (LI-COR).

### 2.3. Candidate Gene Sequencing and Sequence Analysis

The *E. urophylla* genome has not been sequenced yet and at the time of the experiments, the *E. grandis* W. Hill ex Maiden sequence was not released. In that context, primer design was based on gene sequences from other species found in databases. Primers were designed in order to get the whole sequence of the candidate genes ([Table S1](#)). Details of primer design for each gene are given in Supplementary Online Material. PCRs were performed in 10  $\mu$ L reaction mixtures with 10 mM incubation buffer 1X CORE Kit Q-Biogene (Tampon CORE Kit Q-Biogene, MP Biomedical), 0.30  $\mu$ M forward and reverse primers, 1.2 units of *Taq* DNA polymerase (Invitrogen) and 15 ng of template DNA. We used the following PCR program for the amplification: 94°C for 4 min; then 35 cycles (40 cycles for *CesAs*) of 94°C for 30 sec, an annealing step for 1 min at the primer's optimized annealing temperature for 1 min and 72°C for 1 min (2 min for *C4Hs*), followed by a final extension at 72°C for 5 min. PCR products were loaded on agarose gels for electrophoresis and the quantity of PCR product was estimated by image analysis with the freeware Image J v1.45b (Wayne Rasband, NIH). One hundred ng of primer product were sent to High-Throughput Genomics Unit (HTGU, Department of Genome Sciences, Univ. of Washington, Seattle) for direct sequencing. For each gene, the overlapping sequences obtained after direct PCR sequencing were assembled and aligned with Codon Code Aligner software (Licor. 2002-2007 ver. 3.5.2), and manually edited. For heterozygote sequences due the presence of INDELs, the sequences of the two alleles were either deducted manually or with INDELLIGENT software [39].

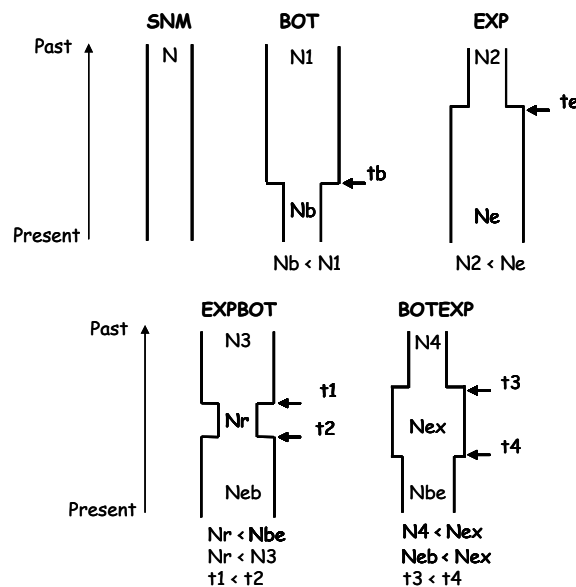
## 2.4. Genetic Structure Analysis

While Tripiiana *et al.* [22] found no population structure on a *E. urophylla* sample encompassing our studied area, we verified the absence of any cryptic population structure on our 43 individual SSR data set by using Structure software (version 2.3.3) [40] [41]. We used sample group information [41] and an admixture model with correlated allele frequencies. 12 independent runs were performed with 100,000 MCMC repetitions and a 100,000 burn-in period for  $K = 1$  to  $K = 4$ . The most likely number of clusters was determined using the log probability of data.

## 2.5. Determining the Demographic History of *E. urophylla* in Timor Island Using Microsatellites Data

We used the Approximate Bayesian Computation (ABC) approach as implemented in DIYABC software [42], [43] to determine the demographic scenario that may have shaped *E. urophylla* diversity pattern observed on SSR data. ABC uses coalescence tool to generate thousands of simulated data having a similar configuration then observed one in terms of individual number, marker type and number and relies on summary statistics to summarize and compare simulated and observed data. Simulations are launched from a set of user-defined demographic scenarios with parameters sampled from prior distributions. DIYABC software provides a full set of tools that we used to simulate SSR data and to perform scenario choice, parameter evaluation and several steps of model validation.

Based on Tripiiana *et al.* [22] results, we considered a single panmictic population and investigated five simple scenarios accordingly: SNM Standard Neutral Model (constant population size), BOT: bottleneck, EXP: demographic expansion, EXPBOT: recent expansion and past bottleneck and BOTEXP: recent bottleneck and past expansion. Cycles of population bottlenecks and expansions are possible in *E. urophylla* due to the volcanic characteristics of Sunda Islands and motivate the exploration of scenario EXPBOT and BOTEXP. These five scenarios are detailed in Figure 2, along with parameter description and imposed constraints. Since we have only scarce information about *E. urophylla* tree, we gave the five scenarios equal prior probability and used non-informative parameter priors, *i.e.* sampled from uniform distribution (Table S2). For each of these five scenarios, we sampled 100,000 simulated data sets whose corresponding information (parameters and summary statistics) was stored in the so-called reference table used in all further analysis except model checking. We recorded mean allele number, mean genetic diversity [44], mean allele size variance and mean Garza-Williamson's  $M$  [45] as summary statistics. Scenario posterior probabilities were calculated using a local logistic regression procedure proceeding on the 10% closest simulated points and that, shortly saying, gives more weight to those closer to the observed data. We then performed a model checking step to verify if the scenario with highest probability fits properly the observed data. As advised by Cornuet *et al.* [43] who recommended not to use for model checking the same summary statistics as for model choice, we constructed a new reference table with same parameter priors than previously but with recording only mean allele number and mean allele size variance as summary statistics and used mean genetic diversity [44] and mean Garza-Williamson's  $M$  [45] for model checking. We simulated then 1000 additional data sets using the parameter posteriors of the scenario with highest probability and compared their summary statistic with those of the observed data. The scenario with highest probability is considered as suitable if the summary statistics of the observed data are included in the 95% confidence interval of the summary statistic's posterior predictive distribution. Parameter posterior distributions were estimated only for the scenario with highest probability using the 10% closest simulated points with a local regression procedure and a logit transformation of the parameters. Confidence in scenario choice was finally evaluated by calculating power and alpha risk associated with choosing the scenario with highest probability. Alpha risk was evaluated by generating 1000 additional simulated sets for each of the four rejected scenario using their respective prior and by assessing the proportion wrongly attributed to the scenario with highest probability based on the reference table. Power was calculated by generating 1000 additional simulated data sets for the scenario with highest probability and by calculating the proportion correctly assigned using the reference table. Confidence in parameter estimates was evaluated by comparing the known parameters of a set of 1000 newly simulated data set obtained for the most likely scenario to those estimated using the current reference table. As measure of parameter estimation error and bias, we calculated the average relative bias and the factor 2, *i.e.* the proportion of pseudo observed data set for which the point estimate is at least half and at most twice the true value [42].



**Figure 2.** Demographic scenarios investigated with ABC analysis. Events are considered backward in time, from present to past. Constraints on the parameters are indicated along with each scenario along with the detail of parameters used with DIYABC. Scenario “SNM” (Standart Neutral Model) considers a population of constant size  $N$ . Scenario “BOT” considers a population of current size  $N_b$  having experienced a sudden bottleneck event  $t_b$  generations ago, from ancestral population size  $N_1$ . Scenario “EXP” considers a population of current size  $N_e$  having experienced a sudden expansion event  $T_e$  generations ago from ancestral population size  $N_2$ . Scenario EXPBOT considers a population of current size  $N_{eb}$  having experienced a recent expansion  $t_2$  generations ago and an ancient bottleneck  $t_1$  generations ago from  $N_4$ . Population size between expansion and bottleneck is denoted  $N_r$ . Scenario BOTEXP considers a population of current size  $N_{eb}$  having experienced a recent bottleneck  $t_4$  generations ago and an ancient bottleneck  $t_3$  generations ago from an ancestral population size of  $N_4$ . Population size between bottleneck and expansion is denoted as  $N_{ex}$ .

## 2.6. Nucleotide Diversity Analysis on Candidate Genes

The SNPs and INDELs were identified by comparison of the 18 genotypes. Polymorphism estimation and neutrality tests were performed with DNAsp software [46]. For all the analyses, the missing data and the INDELs were excluded in all the subsets. Nucleotide diversity on a base pair basis was estimated by  $\theta_w$  from the number of polymorphic segregating (S) sites [47] and by  $\pi$  [44] from the number of pairwise mismatches. These two site frequency spectrum related statistics are two estimators of  $4N\mu$ , with  $N$  effective population size and  $\mu$  the mutation rate per nucleotide and per generation. They are complementary since  $\theta_w$  is not sensitive to allelic frequencies and  $\pi$  is. To determine a possible non-equilibrium status of the studied genes, we measured Tajima’s D [48]. Tajima’s D quantifies the difference between  $\pi$  and  $\theta_w$ . A significant negative value of Tajima’s D can indicate an excess of rare variants as expected under positive selection. It can also reflect demographic event such as population expansion. A significant positive Tajima’s D value, at the opposite, indicates an excess of high frequency variants as expected under balancing selection or diversifying selection. It can also reflect population structure or bottleneck events. We calculated the rate of synonymous (dS) and non-synonymous (dN) substitution according to Nei’s [44] method. The dN/dS ratio should provide insights into the long-term selective pressures on a full gene or a particular gene region and allows us to identify purifying selection (dN/dS < 1) and positive selection (dN/dS > 1). As the later criteria for positive selection is overly stringent because positive selection often only acts on functional domain [49] we also performed a sliding window analysis with win-



dow length of 100 bp and step of 25 bp. We finally performed Hudson Kreitman and Aguade (HKA) test [50] and McDonald & Kreitman (MDK) test [51] with publicly available sequences of *E. pillularis* Smith for *CesA1* (NCBI sequence AB591266.1), *CesA3* (NCBI sequence AB591273.1) and *CAD2* (NCBI sequence AB591254.1) genes as outgroup sequences. HKA and MDK neutrality tests are based on polymorphism to divergence comparison and are much less sensitive to demographic history than neutrality tests based on site frequency spectrum.

## 2.7. Determining Selective Status of Candidate Genes Accounting for Demographic History

Demographic history strongly impact sequence polymorphism pattern and it is necessary to account for it while searching for selection imprint on candidate genes. To achieve that aim, we build the expected null distribution of several summary statistics for sequence data, conditional to the most likely demographic scenario inferred from the SSR data set, and used it to assess neutrality status of the candidate genes. We built this conditional distribution and detected outlier sequences using EGGLIB Python/C++ library and its ABC commands [52]. This library provides tools to easily perform coalescent simulations from inferred parameter posteriors and determine outlier status of candidate loci. Since DIYABC and EGGLIB do not use the same scenario parameters and that SSR and SNPs have different mutation rate and pattern, it is not possible to use the parameter posterior distributions inferred from SSR with DIYABC in our candidate gene analysis. We thus first calibrate scenario parameters for sequence data by running a new ABC analysis using EGGLIB, with candidate gene sequences as observed data and investigating only the most likely scenario inferred from SSR data. We simulated  $10^6$  data sets, using uninformative priors and accounting for gene recombination. As summary statistics, we recorded average  $\Theta_w$  and the relative frequency of 4 classes of minor allele frequency *i.e.* overall proportion of all polymorphic sites from all loci with minor allele frequency: below 0.125, between 0.125 and 0.25, between 0.25 and 0.375 and under 0.375. Parameter posterior distribution was established through simple rejection procedure [53], keeping the 1% simulated data points closest to the observed data and applying a log transformation. To further proceed with posterior based coalescent simulations, we transformed the obtained continuous parameter posterior distributions into a single discrete multivariate distribution. We used the discrete multivariate posterior distribution of the parameters to proceed with two model checking procedures. As the studied candidate genes have different length, we simulated 10,000 times a 500 bp fragment from the posterior parameter distribution to be compared with the set of 500 bp long non overlapping fragments obtained through sliding window procedure on our candidate loci. We first applied a procedure similar to that used by DIYABC, by positioning the mean  $\pi$ ,  $\Theta_w$  and Tajima's D calculated over the 44 observed 500 bp long non overlapping fragment in the corresponding posterior predictive distribution. We then compared the posterior predictive and observed distributions of  $\pi$ ,  $\Theta_w$  and Tajima's D using Kolmogorov-Smirnov [54] test. We finally performed outlier detection by identifying the 500 bp observed fragments with Tajima's D outside the 95% bilateral confidence interval of the posterior predictive distribution. Such outliers were considered as presenting evidence for selection signature.

## 3. Results

### 3.1. Candidate Gene Primary Structure

Each gene consensus sequence was built by alignment of the overlapping fragments. The consensus sequence was then compared with the most similar gene used for the primer design in order to predict the 5' and 3' UTR, as well as the exonic and intronic regions. Hundred percent of the *E. urophylla* *CAD2* gene (*EuCAD2*) was sequenced with a total length of 5408 bp, containing about 2500 bp of promoter region (GQ387647). We found two *EuC4H* copies and hundred percent of both *EuC4H1* and *EuC4H2* genes were sequenced. The sequenced genomic DNA length was 5020 bp for *EuC4H1* (JX270996) and 2850 bp for *EuC4H2* (JX270997). The two sequences were aligned with the three *C4Hs* of *Populus trichocarpa* Torr & A. Gray ex Hook and the two *C4Hs* of *Populus tremuloides* Michx to identify the localization of the exons and introns which have standard GT/AG splicing sites. A dramatic difference between the two *EuC4Hs* primary structures was observed: *EuC4H1* contains 3 exons (785 pb, 134 pb and 599 pb) and 2 introns (676 pb, 1294 pb), while *EuC4H2* had 2 exons (866 pb, 745 pb) and 1 intron (105 pb). Actually, the sequence of the *EuC4H2* exon2 was highly similar to the exon 2 and the exon 3 of *EuC4H1*. At the nucleotide level, the similarity between the whole genes *EuC4H1* and *EuC4H2* is only 25%. However, when we compare the coding sequences, there is 62% similarity. The length of the se-

quenced *EuCesA1* (JX270998), *EuCesA2* (JX70999) and *EuCesA3* (JX271000) genes are, respectively, 6723 bp, 5529 bp and 5493 bp. *EuCesA1* and *EuCesA3* have 13 exons while *EuCesA2* has 12 exons. When we compare the mRNA nucleotide sequence for each copy pair, we obtain 62% to 64% of similarity. As for the whole sequence of each gene, we get 41% to 46% of similarity.

### 3.2. Nucleotide Diversity of Candidate Genes

Each gene was sequenced on 18 individuals (Table 1). Sequences are available from GenBank under accession nos. KF970937- KF971152. On the full sample, the number of sites without missing data and gaps varied from 1027 bp (*EuC4H2*) to 4615 (*EuCAD2*), with number of segregating sites  $S$  varying from 39 (*EuC4H2*) to 170 (*EuCAD2*). Taking all genes together, we had 19,798 sites without any gap or missing data, cumulating 661 segregating sites. The overall level of diversity was 0.00651 for  $\pi$  and 0.00744 for  $\theta_w$ . However, at a gene level, the diversity varied on a twofold extends: for  $\pi$ , from 0.00479 (*EuCesA1*) to 0.00899 (*EuC4H2*) and for  $\theta_w$ , from 0.0051 (*EuCesA3*) to 0.00916 (*EuC4H2*). For the six studied genes,  $dN/dS$  ratio always stayed below one at the full length scale, varying on almost a ten fold extend, from 0.0167 for *EuCesA3* to 0.1341 for *EuCesA2*. Considering the sliding window analysis with window length of 100 bp and step of 25 bp, 3 gene regions exhibited  $dN/dS$  ratio over 1:1.786 between 1360 and 1509 bp in *EuCesA2* (beginning of exon 5), 1.029 between 3363 and 3462 bp in *EuCesA3* (exon 9) and 2.73 between 5343 and 5442 bp in *EuCesA1* (exon12). Tajima's  $D$  values measured at whole gene scale varied from  $-1.05718$  (*EuC4H1*) to  $0.31332$  (*EuCesA3*) and none of them were significantly different from zero according to standard test procedure based on Wright Fisher equilibrium hypothesis. Summary statistics recorded on each gene of the full sample are presented in Table 2. Table 3 presents detailed value of  $\pi$ ,  $\theta_w$  and Tajima's  $D$  for fragment defined in the 500 bp non overlapping fragment set. Note that three of these fragments have no polymorphism at all. Neither HKA nor MDK test results indicated significant deviation from neutral expectation for the evaluated genes (*EuCesA1*, *EuCesA3* and *EuCad2*).

### 3.3. Genetic Structure Analysis on SSR Data

Using Structure, we found no cryptic population structure. Increasing group number from 1 to 4 did not increase likelihood model. Average likelihood for  $K = 1$  was  $-3727$ , average likelihood for  $K = 2$  was  $-3764$ , average likelihood for  $K = 3$  was  $-3822$  and average likelihood for  $K = 4$  was  $-4122$  with low standard error between repetition of  $K$  (from 3.28 for  $K = 1$  to 384.05 for  $K = 1$ ). As an additional proof of no population structure, examination of individual repartition among group for  $K > 1$  shows high admixture level for individual with minor group contribution.

### 3.4. Demographic History of *E. urophylla* in Timor Island Based on Microsatellite Data

The reference table was built with a total of 513,800 simulated data points, evenly distributed across scenarios: 102,500 for scenario SNM, 102,361 for scenario BOT, 102,951 for scenario EXP, 102,797 for scenario EXPBOT and 103,191 for scenario BOTEXP. Logistic approach indicated the highest posterior probability for scenario EXP with a mean value of 0.527 and confidence interval [0.502; 0.551]. BOTEXP and EXPBOT had lower probability, respectively 0.311 with confidence interval [0.287, 0.334] for BOTEXP and 0.134 with con-

**Table 2.** Diversity pattern on candidate genes.

Gene	$l$	$S$	$\pi$	$\theta$	$D_{taj}$	$dN/dS$
<i>EuCesA1</i>	4099	108	0.0048	0.0064	$-0.9171$	0.0919
<i>EuCesA2</i>	4427	149	0.0073	0.0081	$-0.3717$	0.1341
<i>EuCesA3</i>	4063	86	0.0055	0.0051	0.3133	0.0167
<i>EuCAD2</i>	4615	170	0.0076	0.0089	$-0.5404$	0.0570
<i>EuC4H1</i>	1567	59	0.0065	0.0091	$-1.0572$	0.1207
<i>EuC4H2</i>	1027	39	0.0090	0.0092	$-0.0645$	0.0217

$l$ : length in base pair, excluding gap and missing data,  $S$ : number of segregating sites,  $\pi$ :  $\pi$ ,  $\theta_w$ :  $\theta$  defined by Watterson,  $D_{taj}$ : Tajima's  $D$ ,  $dN/dS$ : ratio of non synonymous to synonymous polymorphism.



**Table 3.** Description and summary statistics for the set of 500 bp non overlapping fragment.

Gene	Firstbase	Lastbase	$\pi$	$\theta_w$	Dtaj	Sim <i>P</i> value
CESA1	1	1400	0.0033	0.0072	-1.7712	
CESA1	1401	2561	0.0038	0.0058	-1.0778	
CESA1	2562	3175	0.0087	0.0072	0.6363	
CESA1	3176	4452	0.0035	0.0068	-1.5630	
CESA1	4453	5119	0.0045	0.0063	-0.9042	
CESA1	5120	5619	0.0040	0.0058	-0.9812	
CESA1	5620	6121	0.0037	0.0068	-1.4692	
CESA1	6122	6624	0.0054	0.0043	0.7353	
CESA1	6625	6723	0.0126	0.0097	0.0871	
CESA2	1	922	0.0187	0.0125	1.7004	*
CESA2	923	1484	0.0153	0.0116	1.1174	
CESA2	1485	2030	0.0060	0.0077	-0.7469	
CESA2	2031	2718	0.0033	0.0072	-1.7654	
CESA2	2719	3220	0.0038	0.0048	0.6462	
CESA2	3221	4098	0.0045	0.0087	-1.6022	
CESA2	4099	4602	0.0024	0.0029	-0.4783	
CESA2	4603	5102	0.0056	0.0068	-0.5568	
CESA2	5103	5529	0.0060	0.0113	-1.5682	
CESA3	1	533	0.0037	0.0077	-1.7019	
CESA3	534	1783	0.0048	0.0048	-0.0105	
CESA3	1784	2607	0.0102	0.0077	1.0550	
CESA3	2608	3107	0.0077	0.0063	0.7413	
CESA3	3108	3607	0.0056	0.0053	0.1813	
CESA3	3608	4235	0.0022	0.0015	1.1066	
CESA3	4236	5132	0.0064	0.0043	1.3949	*
CESA3	5133	5633	0.0044	0.0039	0.4069	
CESA3	5634	5712	0.0000	0.0000	NA	
CAD2	1	555	0.0050	0.0058	-0.4140	
CAD2	556	1064	0.0071	0.0063	0.4061	
CAD2	1065	1586	0.0074	0.0072	0.0943	
CAD2	1587	2164	0.0109	0.0101	0.2607	
CAD2	2165	2681	0.0069	0.0121	-1.4819	
CAD2	2682	3208	0.0093	0.0106	-0.4372	
CAD2	3209	3764	0.0085	0.0097	-0.4063	
CAD2	3765	4747	0.0080	0.0116	-1.0500	
CAD2	4748	5249	0.0047	0.0048	-0.0827	
CAD2	5250	5408	0.0106	0.0168	-1.0831	

## Continued

C4H1	1	2202	0.0036	0.0072	-1.6328
C4H1	2203	3198	0.0077	0.0111	-1.0413
C4H1	3199	4907	0.0089	0.0101	-0.4123
C4H1	4908	5021	0.0000	0.0000	NA
C4H2	1	560	0.0148	0.0150	-0.0266
C4H2	561	1374	0.0036	0.0039	-0.1713
C4H2	1375	2850	0.0000	0.0000	NA

“Firstbase” and “lastbase” define the fragment boundaries;  $\pi$ :  $\pi$  (ref)  $\theta_w$ :  $\theta$  defined by Watterson (ref),  $D_{Taj}$ : Tajima’s D; “Sim  $P$  value” indicated significant  $P$  values for Tajima’s D calculated using Tajima’s D expectation under sudden expansion demographic history and no selection, \*:  $0.025 > P$  and  $P > 0.975$ , \*\*:  $0.005 > P$  and  $P > 0.995$ .

fidence interval [0.119, 0.149] for EXPBOT. The two remaining scenarios have considerably lower probability, below 1% for scenario BOT and below 5% for scenario SNM. We kept only the scenario EXP for further analysis. Using the new reference table constructed for model checking and using only 2 summary statistics, scenario EXP still appears to be the most likely scenario. Model checking procedure further indicated that scenario EXP is compatible with observed data with a  $P(\text{simulated} < \text{observed})$  of 0.265 for mean genetic diversity and of 0.403 for mean Garza-Williamson’s  $M$ . On the contrary, scenario BOTEXP, the second scenario in terms of posterior probability, was not compatible with observed data as  $P(\text{simulated} < \text{observed})$  for mean genetic diversity was 0.976. Main characteristics of parameter posterior distributions for scenario EXP are presented in [Table 4](#). For most parameters, the 95% confidence interval appeared to be broad, but examination of the posterior distributions indicated that they generally markedly differed from the priors ([Figure S1](#)). The posterior distribution of  $N_2/N_e$  indicates a marked expansion (5% confidence interval [0.001; 0.110]), with a median value of 0.014. Confidence in parameter estimate is good ([Table 4](#)), with a mean bias below 1, except for  $N_2$  (mean bias 1.448) and Fact 2 above 0.7 (except again for  $N_2$ , Fact 2 = 0.634). These values indicate that parameter estimation is reasonably good most parameters except  $N_2$ . We finally assessed confidence in scenario choice by calculating associated power and alpha risk. The power to identify scenario EXP is good (81.5%) and its alpha risk is moderate (20.35%).

### 3.5. Determining Selective Status of Candidate Genes Accounting for Demographic History

Following results obtained from ABC analysis conducted on SSR data, we build the expected null distribution of several summary statistics for sequence data conditional to the EXP scenario and used it to assessed neutrality status of the candidate genes. We first evaluated the EXP scenario parameters using candidate gene data. Main characteristics of parameter prior and posterior distributions for scenario EXP are presented in [Table 5](#) and they are graphically displayed in [Figure S2](#). Posterior median value for the ratio of ancestral size to actual size is 0.395, with corresponding 95% being [0.098; 0.778]. Median posterior value for current population size ( $4\mu N_0$ ) was 0.011 with a narrow confidence interval [0.007; 0.025]. For expansion time expressed in  $4N_0$  generations, median value was 0.273 with broad confidence interval ([0.026; 0.480]).

The parameter posterior distributions obtained for scenario EXP were further used to simulate ten thousand 500 bp long sequence fragments. Tajima’s D,  $\theta_w$  and  $\pi$  values were calculated on each simulated fragment and used to determine a 95% bilateral confident interval characterizing the expected values of these statistics under a neutral model. The two model checking procedures indicated that observed data are consistent with EXP demographic model and inferred parameters. Positioning Tajima’s D values calculated on the 500 bp non-overlapping observed fragment set allow identifying two fragments presenting outlying values as indicated in [Table 3](#). These two fragments, located in *EuCesA2* and *EuCesA3* are positioned on [Figure 3](#) along with gene exonic structure,  $dN/dS$  and Tajima’s D sliding window analysis and SNPs associated with wood traits. The first fragment is positioned in *EuCesA3*, between base 4236 and 5132 and has a positive Tajima’s D (1.395,  $P$  value < 0.025). The second outstanding fragment is positioned in *EuCesA2*, between bases 1 to 922 and has also a positive Tajima’s D value (1.700,  $P$  < 0.025) Note that this *EuCesA2* fragment contains 2 SNPs found to be associated with Acid Soluble Lignin by Denis *et al.* [17].

**Table 4.** Parameter estimates for scenario EXP (demographic expansion) using the Approximate Bayesian Computation approach (Cornuet *et al.* 2008, 2010).

Parameter	Mean	Median	IC95	Mean bias	Fact2
N2	3 659	2 828	[345; 11,140]	1.448	0.634
Te	3 703	3 091	[333; 9458]	0.594	0.772
Ne	2.5E+05	2.4E+05	[2.9E+04; 4.9E+05]	0.619	0.738
Mean_μmic	2.7E−04	2.0E−04	[1.0E−04; 8.4E−04]	−0.058	0.797
Mean_P	2.199	1.926	[0.520; 4.764]	0.115	0.756

IC 95% indicates 95% confidence interval; MeanBias indicates the average relative bias and Fact2 the average proportion of pseudo observed data set for which the point estimate is at least half and at most twice the true value; N2: ancestral effective population size; Ne: current effective population size; Te: time of expansion in generation number; Mean\_μmic: mean mutation rate; Mean\_P is average parameter of the Generalized Stepwise model geometric distribution.

**Table 5.** Prior specifications and posterior estimates for demographic parameter of scenario 3 as inferred from 6 candidate genes using ABC approach (De Mita & Sio 2011).

	Priors			Posteriors		
	Dist	Min	Max	Mean	Median	IC 95
THETA	U	0	0.03	0.012	0.011	[0.007; 0.026]
ANCSIZE	U	0	1	0.406	0.395	[0.098; 0.778]
DUR	U	0	0.5	0.137	0.097	[0.009; 0.410]
RHO	U	0	0.03	0.011	0.01	[0.001; 0.026]

THETA defines actual population size  $\times$  mutation rate, ANCSIZE is the ratio of ancestral population size to actual population size and DUR indicates the duration of expansion. For each of these three parameters, prior distribution shape (U: uniform) is given with minimum (min) and maximum (max) values. Posterior mean, median and 95% confidence intervals (IC95) are also presented.

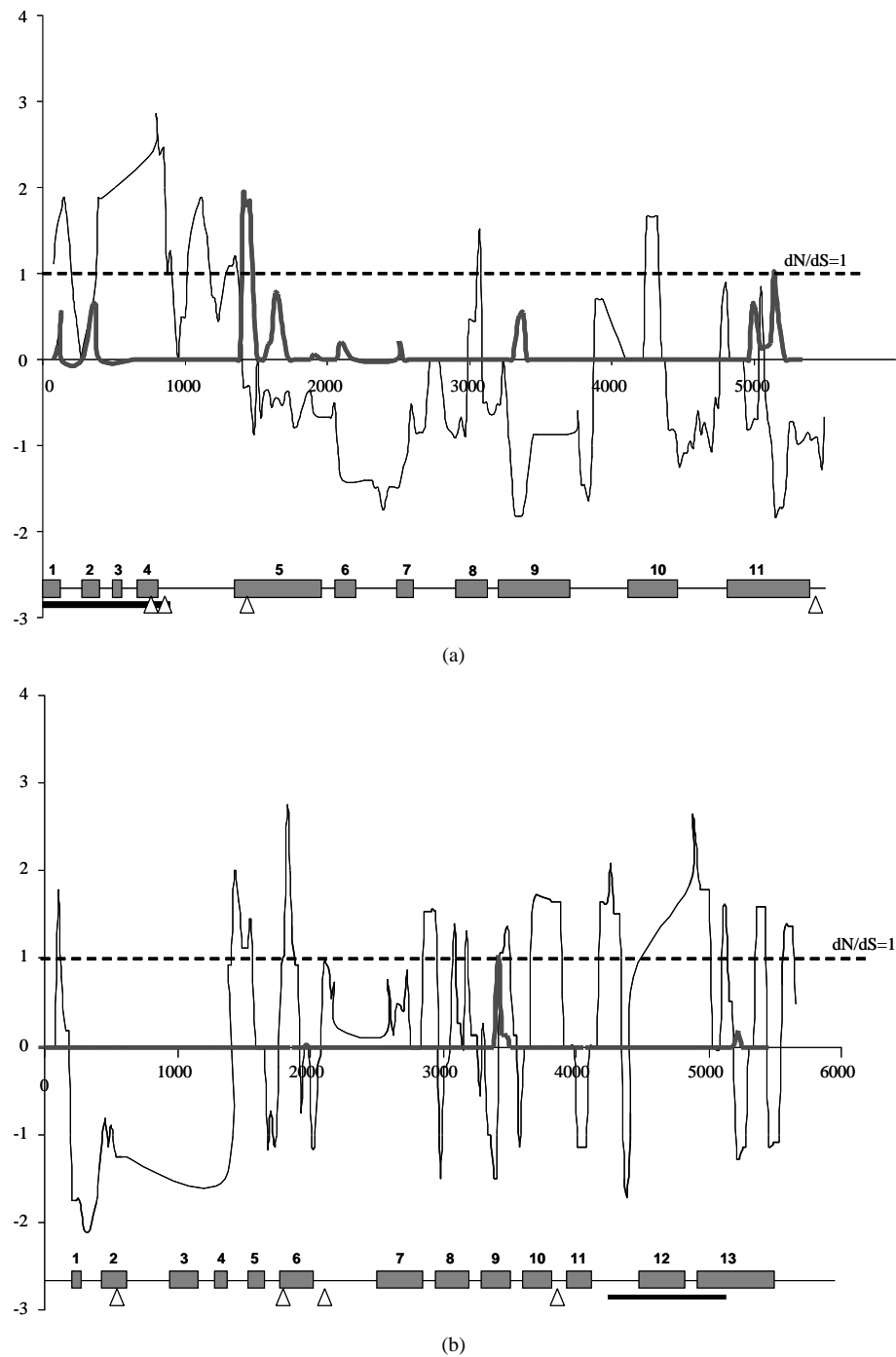
## 4. Discussion

We studied the diversity pattern of six full length candidate genes involved in wood formation in *E. urophylla*, a tree originating from the Sunda Island and presenting high diversity in various phenotypic traits. This work brings original results to understand impact of natural selection and demography on natural populations for genes coding for crucial traits in trees.

### 4.1. Genetic Structure, Demographic History of *E. urophylla* and Consequence on Polymorphism Pattern

*E. urophylla* is among the rare eucalypts species not originating from Australia. It was formerly found to be not structured at the nuclear level [22] [31] and it is believed to have undergone several cycles of population expansion and bottleneck in relationship with the volcanic nature of Sunda Islands [31]. Our microsatellite data analysis based on a tree sample from seeds originating from a small region of Timor Island is consistent with the absence of genetic structure and with a sudden demographic expansion. If local extinctions due to volcanic eruption may have occurred, it didn't affect markedly overall diversity pattern in the sampled region where the *E. urophylla* populations are known to be large and stable. This expansion is confirmed by the ABC analysis we performed on gene sequence data set alone (results not shown). Previous studies on *E. urophylla* didn't investigate possible demographic expansion but only tested bottleneck [32]. One of the main consequences of population expansion on polymorphism pattern is an excess of rare variants and thus negatively skewed Tajima's D values for sequence data. In such a situation, not accounting for population demography may lead to the detection of false positive hits for positive directional selection.

The two data sets we used (microsatellites and genes) are very different in terms of sample size and marker mutation rate and modality. However, it is interesting to compare their respective posterior distribution for ance-



**Figure 3.** Sliding window analysis of *EuCesA2* (a) and *EuCesA3* (b) genes with window length of 100 bp and 25 bp steps. Black thin lines indicate Tajima's D value and grey thick lines figure dN/dS ratio. Dashed lines materialize the dN/dS threshold value of one. At the bottom of each graph gene structure was schematized with numbered exons represented by grey boxes. 500 bp long regions with outstanding Tajima's D value are represented by a black horizontal thick line and SNPs found to be associated with wood traits are identified by white triangles (Denis *et al.* 2013)

stral to current population size ratio in ABC expansion scenario. Indeed it does not depend on mutation rate and is a measure of expansion intensity. Expansion appears much stronger in microsatellites data set (median value

of ANCSIZE, the ratio of ancestral population size to actual population size, is 0.014% and 95% confidence interval [0.001; 0.110]) than in gene data set (median value 0.395% and 95% confidence interval [0.098; 0.778]). Two explanations can be proposed for this discrepancy. On the first hand, some uncertainties can be associated with parameter posterior distribution estimation from ABC procedure. For example, in the ABC analysis performed with the microsatellite data set, the mean bias for ancestral size  $N_2$  is 1.448, which is beyond the threshold of 1. Poor parameter posterior estimation is often the result of an observed data set insufficiently informative [42]. On the second hand, our gene data set is built only with wood formation genes that are likely impacted by natural selection. For example, balancing selection may generate intermediate frequency variants that may attenuate demographic expansion signal. Finally, the choice of the proper posterior demographic parameters is very important and critical because it strongly affects false discovery rate. In the particular case of this study, we choose to be conservative and to use the posteriors inferred from the candidates themselves to draw neutral expectations, with in mind the fact that they may not exactly correspond to selective neutrality.

#### 4.2. Selection Signature in Wood Related Genes of *E. urophylla*

We investigated possible natural selection imprint on wood formation candidate genes using several complementary approaches: a test based on site frequency spectrum (Tajima's  $D$ ) with ad-hoc correction for neutral expectations, non-synonymous comparison to synonymous polymorphism ( $dN/dS$  ratios) and two selection tests based on polymorphism to divergence comparison (MDK and HKA tests). As the latter approach required an outgroup sequence, we were not able to implement the two corresponding tests on the full sequence data set.

None of the studied gene exhibited a clear selection imprint on its full length but we could observe interesting trends in global  $dN/dS$  between copies of the same gene family, witnessing possible contrasted constrain level. According to Ohno's [56] gene family classical model, contrasted constrain level can be expected as at least one member of the family should maintain the original function of the gene, and the other copies possibly undergoing pseudogenization or subfunctionalization. The three members of the *CesA* gene family we studied are all involved in secondary cell wall formation and possess homologous counterparts across angiosperms [27]. While these three genes seem to be co-expressed in secondary tissues, *E. grandis CesA3* copy was found to display the highest expression level in Ranik *et al.* [27] study. In this work, the three *CesA* gene copies present very contrasted  $dN/dS$  ratio, with almost a 10-fold ratio between *EuCesA3* (0.0167) and *EuCesA2* (0.1341). While these  $dN/dS$  ratios indicate purifying selection in both cases, they suggest more constraints on *EuCesA3* than in *EuCesA1* and *EuCesA2*. Similarly, *EuC4H2* ( $dN/dS$  ratio of 0.0217) appears to be more constrained than *EuC4H1* (0.1207). The only *C4H* copy of *A. thaliana* is expressed in all tissues and responds to light, wounding and fungal infection [28], indicating that it plays various functions in phenylpropanoid metabolism that open a door for subfunctionalization of gene copies. Indeed citrus sinensis *C4H1*, ortholog of *EuC4H2* is wound inducible unlike *C4H2*, ortholog of *EuC4H1* [30].

Based on site frequency approaches and by focusing on the 500 bp length non-overlapping fragment set, we found two 500 bp regions that could bear the imprint of natural selection. These two regions both present an excess of sites in intermediate frequencies that can be interpreted as balancing selection. The first region is located between bases 4236 and 5232 of *EuCesA3* and covers part of intron 11, exon 12, intron 12 and part of exon 13. This particular region does not itself contain a SNP investigated by Denis *et al.* [17] but a nearby SNP (*EuCesA3*-3854) was found to be associated with cellulose and extractive related traits [17]. However, as we do not have proper outgroup sequence for this *EuCesA3* region, it is not possible to make any final conclusion about its selective status. The second region is located in *EuCesA2* between 1 bp and 922 bp and encompasses the four first exons, the three first introns and part of the fourth exon. It contains two SNPs (*EuCesA2*-0762 and *EuCesA2*-853) found to be associated with acid soluble lignin by Denis *et al.* [17], among them one being non-synonymous. We also do not have a outgroup sequence for this *EuCesA2* particular region but Sexton *et al.* 2011 found an unexpectedly high concentration of trans-subgeneric conserved SNPs in the nearby promoter interpreted as a signal of balancing selection.

Besides our work, several studies reported evidence of balancing selection in *CesA* genes in evolutionary distant tree species: in *Pinus pinaster* (Aiton) [19], *Pinus radiata* D. Don [18], in Eucalyptus genus in general [55] and in *E. urophylla* [56] in particular. This suggests the wide adaptive importance of *CesA* genes across trees and raises the question of functional significance of this result. As pinpointed by some authors, wood formation genes may play an important role in plant defence [57], [58] although the modalities are not clear.



## Authors' Contributions

LCK: data analysis BF, SU DNA extraction and gene study, JP: SSR genotyping, LC-K, BF, SU JMB: manuscript preparation. JMB: funding and overall supervision. All authors read and approved the final manuscript.

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## Appendix

**Table S1.** Primers used for gene amplification.

Gene	Forward primer			Reverse primer			Annealing temperature
	Location	ID	Sequence	Location	ID	Sequence	
EuCesA1	Promoter	EgCesA1_-276.22F	GCTTCAACACA ATGACACCAAC	Exon 2	EgCesA1_212.23R	AATTCATTTCTC TCAGCATCAGC	59
	Exon 1	EgCesA1_108.22F	CAAGGCTTGTG TCGAATATGAG	Exon 4	EgCesA1_421.22R	GAAGCTTGAGCC TCTTTCTCAGC	62
	Exon 3	EgCesA1_275.23F	GACACATCAGC AGTGTCTTCTACG	Exon 5	EgCesA1_693.23R	AGTAATCCTAT TCACGGGAGACC	59
	Exon 5	EgCesA1_629.20F	TTTGTTTCGCAT ACTCATGG	Exon 6	EgCesA1_918.23R	CGTTTCAACAAG AGACTCGAATG	59
	Exon 6	EgCesA1_796.22F	CCTTTGATCACT GCCAATACAG	Exon 9	EgCesA1_1379.22R	CAGTCAAGGTT GAGGATGTACG	59
	Exon 8	EgCesA1_1245.20F	ACTTCCTCGGC TGGTGTATG	Exon 11	EgCesA1_1892.23R	AACAGCGTCGA CTCAATAAACAC	62
	Exon 10	EgCesA1_1636.22F	TCTATGCCGAA TTTACCAAAGC	Exon 12	EgCesA1_2310.23R	AATTGTGCAATA AGCGACAAGAG	59
	Exon 12	EgCesA1_2167.25F	TCCGTGAAAATA TTCTTGAGTAGAC	Exon 13	EgCesA1_2706.21R	ATAACCTTTGT TCAGCGCATC	59
	Exon 13	EgCesA1_2518.22F	GGATTCTGAAG ATGTTAGCTG	Exon 13 (3'UTR)	EgCesA1_3214.20R	ATTCCATGCAT CGCACATTC	59
EuCesA2	Promoter	EgCesA2_-152.22F	TGACGTCTCTTC AACTCAAACC	Exon 2	EgCesA2_144.18R	GTCCTTCAGG CCGATCTC	62
	Exon 2	EgCesA2_124.18F	GACGAGATC GGCCTGAAG	Exon 3	EgCesA2_339.21R	AAACTCGTCC TCGAAATCCTC	62
	Exon 3	EgCesA2_295.19F	GACGAGGAC GACCACTTCG	Exon 5	EgCesA2_597.22R	GTAGTCATCTT CCTCTCCATCG	54
	Exon 5	EgCesA2_495.20F	GGAGTGGA GGAGAGGATCG	Exon 6	EgCesA2_843.23R	AGTTTCTCTGTC TATCGGGTTCC	54
	Exon 6	EgCesA2_735.22F	AACTGATGCA TTCCCTCTATGG	Exon 6	EgCesA2_1184.21R	CTGGGCTCCAC CTTATCTTTC	60
	Exon 6	EgCesA2_1108.20F	TATAGCATCG AGCCGAGGAC	Exon 9	EgCesA2_1615.22R	CATAGCAAAG CTTCTTTCCAAG	54
	Exon 8	EgCesA2_1385.21F	AAGGTAAGG AGTTGCCTCGAC	Exon 10	EgCesA2_2008.23R	AACCCTCAAGT CCTTCTCAATG	57
	Exon 10	EgCesA2_1878.23F	GAAGGATGAT ACGAGTTTGCTTG	Exon 11	EgCesA2_2488.23R	TGGAAGTGAAA GGGTAGACAATG	60
	Exon 11	EgCesA2_2399.22F	ATTGTCCTTTG TGGTATGCTTG	Exon 12	EgCesA2_2917.23R	ACGATCCGTAT CCATTGTTTATG	60

## Continued

	Exon 12	EgCesA2_2714.20F	TATTCCAAGG CCTCCTCAAG	Exon 12 (3'UTR)	EgCesA2_3315.23R	TTTCAGTTCCA CACTCTTCAAGC	60
EuCesA3	Promoter	EgCesA3_-194.21F	AGCTTAGCTTC AAGGCAATGG	Exon 1	EgCesA3_189.21R	CAAATCTCCAT CAACCGTGAG	65
	Exon 1	EgCesA3_51.19F	ACTTGTCGCC GGTCTCAC	Exon 2	EgCesA3_303.22R	TTTGAGACGCT TGTATCGAGTC	67
	Exon 2	EgCesA3_246.20F	GAGGAGAGAA GGGAGCCAGT	Exon 3	EgCesA3_362.20R	TCGTGCTCGAG ATCATCAAT	65
	Exon 3	EgCesA3_317.20F	TGGAGGGTGA TGATGATGAA	Exon 4	EgCesA3_578.21R	GGATGAACTCG TTTGTGAAGG	63
	Exon 4	EgCesA3_526.22F	CCTTACCACTC CTTGCTACTG	Exon 5	EgCesA3_713.19R	CTTCGTCATCA GACGCTTTG	65
	Exon 5	EgCesA3_629.21F	TGTGTCAGCT CACTTGTTTGC	Exon 6	EgCesA3_975.21R	GTCTTTAGCA CGAACGGATCA	65
	Exon 6	EgCesA3_866.23F	GATGGGGAG ACAGAACCGTA	Exon 7	EgCesA3_1135.22R	CGGAAGAAAA CGCTTATAGCC	63
	Exon 7	EgCesA3_994.21F	TCATCTTATGG TCACGGAGAGA	Exon 8	EgCesA3_1496.20R	ATGTCCGGGT CATTGATGT	59
	Exon 8	EgCesA3_1439.21F	GAGGGTGGAA AGAAAGAATGG	Exon 10	EgCesA3_1932.23R	GCGATCAAGA TAGGTCTCACG	65
	Exon 10	EgCesA3_1872.21F	TTGGTCTTTGGT TAACATCCATC	Exon 11	EgCesA3_2156.22R	AGCAGGAAATC TTATCGACTGG	65
	Exon 11	EgCesA3_2041.21F	AGGGAAGGTGA ACCCAATATG	Exon 12	EgCesA3_2539.21R	AGACCTCCGCT GTGTCCAAG	65
	Exon 12	EgCesA3_3009.20F	ACAACGCAAA GGATCATCCAG	Exon 13	EgCesA3_3392.21R	GACCATCTTAG GACGCTTAGGAC	65
	Exon 13	EgCesA3_2709.21F	CCGGAGACAG GCATTATATGG	Exon 13	EgCesA3_3116.21R	AGTGCTGCTGG ACTTGAAGAAG	65
	Exon 13	EgCesA3_2504.22F	GAGCTGTTGAT GTCCGAGATG	Exon 13 (3' UTR)	EgCesA3_2809.20R	TGGCCGGAAGA GTACAGTAGG	65
EuCAD2	Promoter	sCIREu1U	GATCCGCCTC CAGAGATAG	Promoter	sCIREu1L	GCTTCCTGTA CTTGTCCTC	58
	Promoter	sCIREu2U	TGCTTCTTCTC CTGATGAC	Promoter	sCIREu2L	GTTCTTGCTG ATCCACAAC	58
	Promoter	sCIREu3U	TTTGCTGATC TCTCTCTCGG	Promoter	sCIREu3L	ATTCCCTAAA ATCTTCTCTGGC	58
	Promoter	sCIREu4U	AAAAGTAAAC GATTGGACGGAC	Promoter	sCIREu4L	TGTTGGATTTC ATTGGCTTG	54

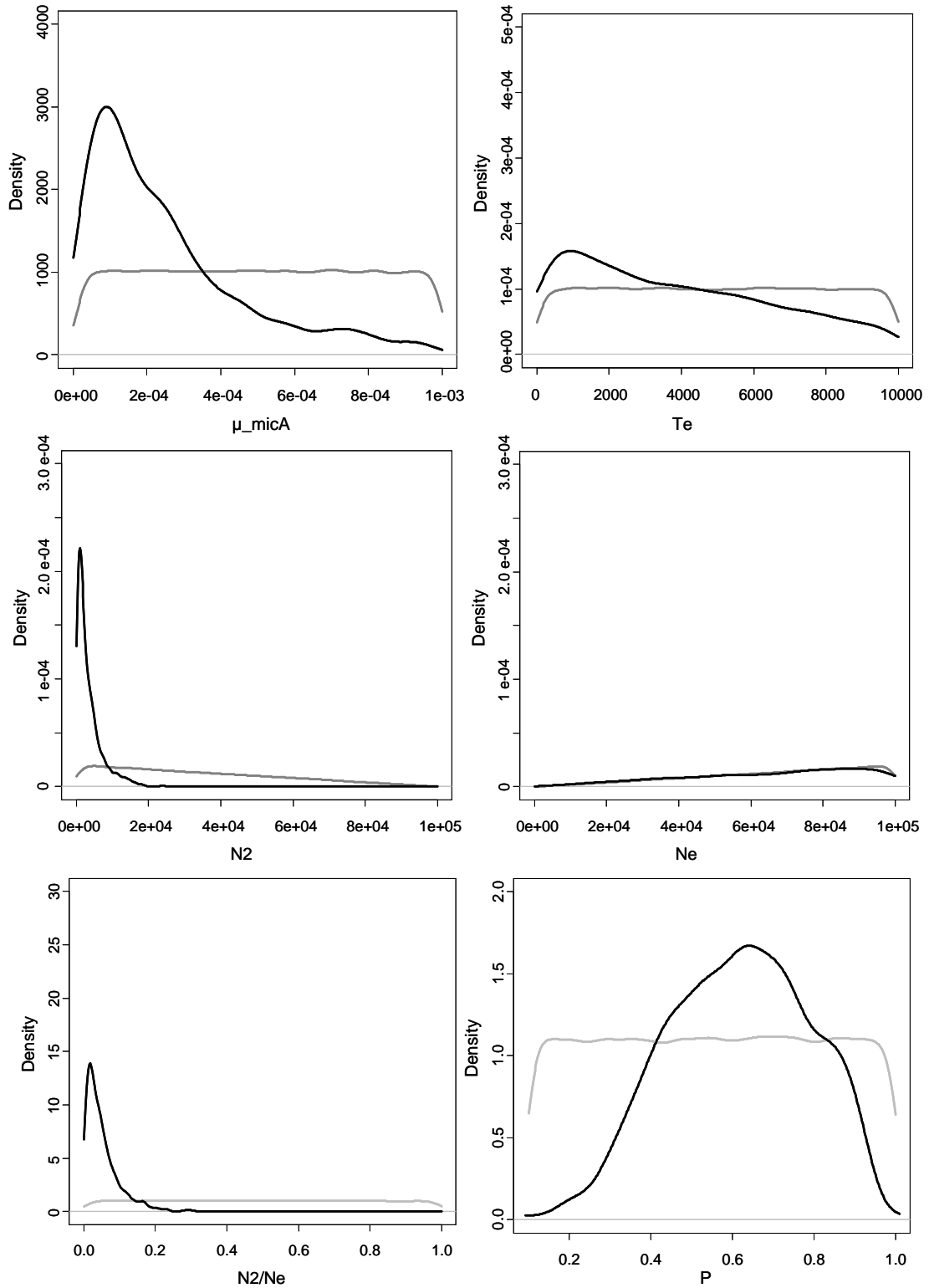


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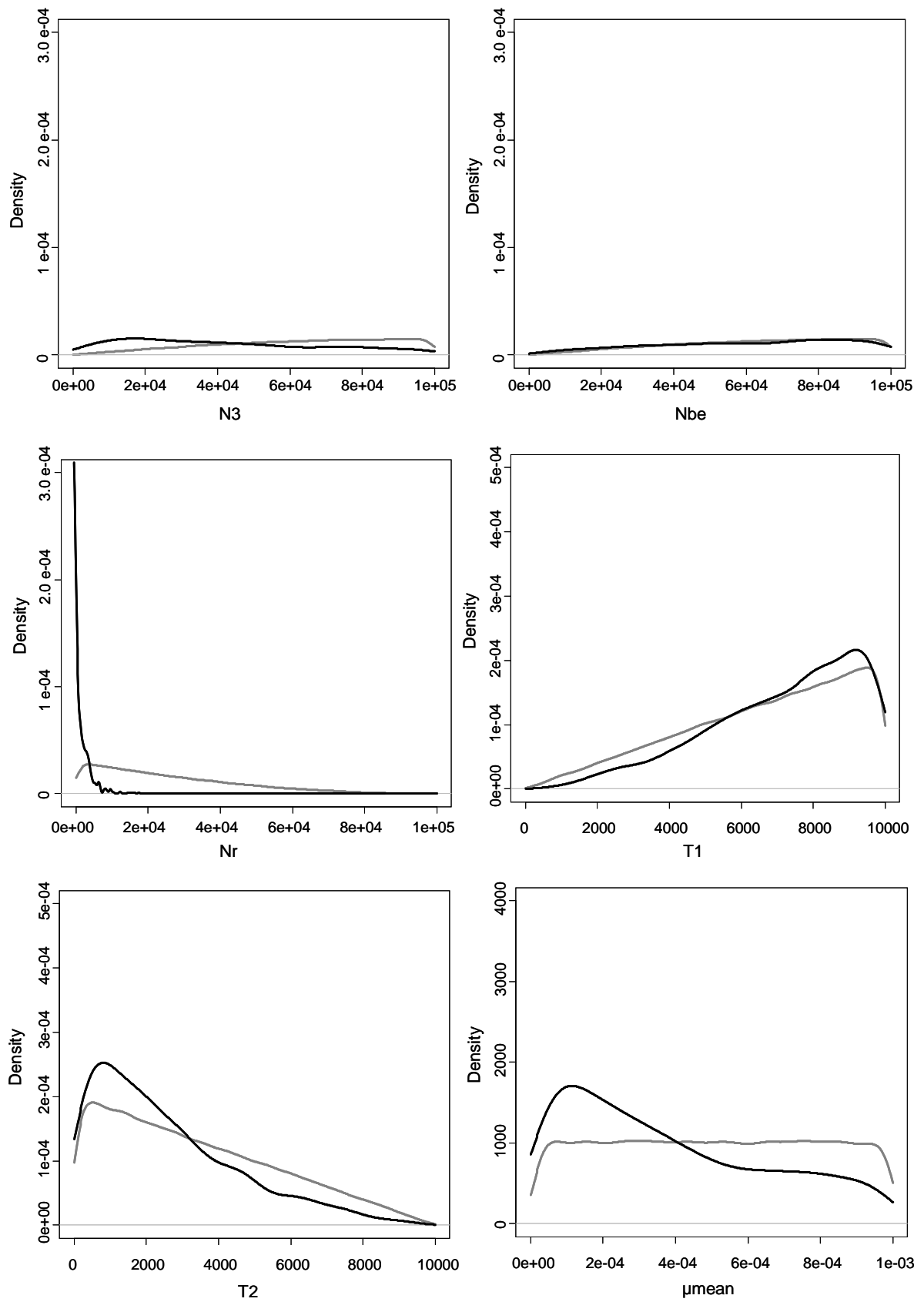
	Promoter	sCIREu5U	GACGAATGGC AAAGCAGAC	Promoter	sCIREu5L	GACACCTCAG CAAGCAATACC	54
	Promoter	sCIREu6U	TGTCAGAAGC ACAGAAACTG	Intron 1	sCIREu6L	ACCTCAGACC AATCAATCG	54
	Exon 1	sCIREu7U	GCCTCAGGTA GATTCAAGAAC	Exon 3	sCIREu7L	CCACTATCTC ACCAGCAAAC	58
	Exon 3	sCIREu8U	GTACACCGA TGGCAAGC	Intron 3	sCIREu8L	CAAACCTGA AGGGCAAAG	58
	Intron 3	sCIREu9U	GTAAACCCGT ACAAAGTGAAAC	Exon 4	sCIREu9L	ATTGAAGAG GAGCATTGATG	58
	Exon 4	sCIREu10U	ACACTATCCC TGTGGTTCAC	Exon 5	sCIREu10L	CCTGACATCA TTCTTCTCG	58
	Exon 5	sCIREu11U	GCAAAGAAA AGGGATTGAC	3' UTR	sCIREu11L	CACTTAGGC AGAAAAGCATC	58
EuC4H1	Promoter	EuC4Hg1-p1U	TTTCCTCTTCC ATCATTTCTC	Promoter	EuC4Hg1-p1L	GAACGCATC TGACTTTGAG	56
	Promoter	EuC4Hg1-p2U	GAAAGAGAGA GGGGTGAATGC	Exon 1	EuC4Hg1-p2L	GAGCCAGTTG CCGAAGATG	60
	Promoter	EuC4H1bisU	AAARCCCTCC GCCTCTGTC	Intron 1	EuC4Hg1-1L	ACGAGATTG CTTACTTCC	54
	Exon 1	EuC4Hg1-2U	GACCCGCTCT TCGTCAAG	Intron 2	EuC4Hg1-2L	GGCTTTGCTA TTTCCCCAG	54
	Intron 1	EuC4Hg1-5U	CCGTGGGAGT TATTGTCTGC	Intron 2	EuC4Hg1-2L	GGCTTTGCTA TTTCCCCAG	58
	Exon 1	EuC4Hg1-2U	GACCCGCTCT TCGTCAAG	Exon 2	EuC4Hg1-2bisL	GATTTCGCCC TTCTGCTG	60
	Intron 2	EuC4Hg1-3U	TGTATAGGTGA AGCGAAAC	3'UTR	EuC4Hg1-3L	TCCAAGTG AAATCAACAG	52
	Intron 2	EuC4Hg1-3U	TGTATAGGTG AAGCGAAAC	Exon 3	EuC4Hg1-6L	ACCAGGATC TTGCTCTCGG	58
	Exon 3	EuC4Hg1-4U	CTGTTGATTT CACTTGGAC	3' UTR	EuC4Hg1-4L	CATAACTACG CCTGTAGACC	54
EuC4H2	Promoter	EuC4Hg2-p1U	CTGATGTGAG ACGGTGTTG	Exon 1	EuC4Hg2-p1L	AGTTGGCGAG AATAATGAGC	58
	Promoter	EuC4Hg2-1C_U	GAAGGTGTGA AGCGAAGATAC	Intron 1	EuC4Hg2-1C_L	CCTGACAATC AAAAGCAAAG	54
	Intron 1	EuC4Hg2-2U	GATTCTCTGT TTTCGTGTTG	3' UTR	EuC4Hg2-2L	GGTTACCACT CCCTTTGAGC	54

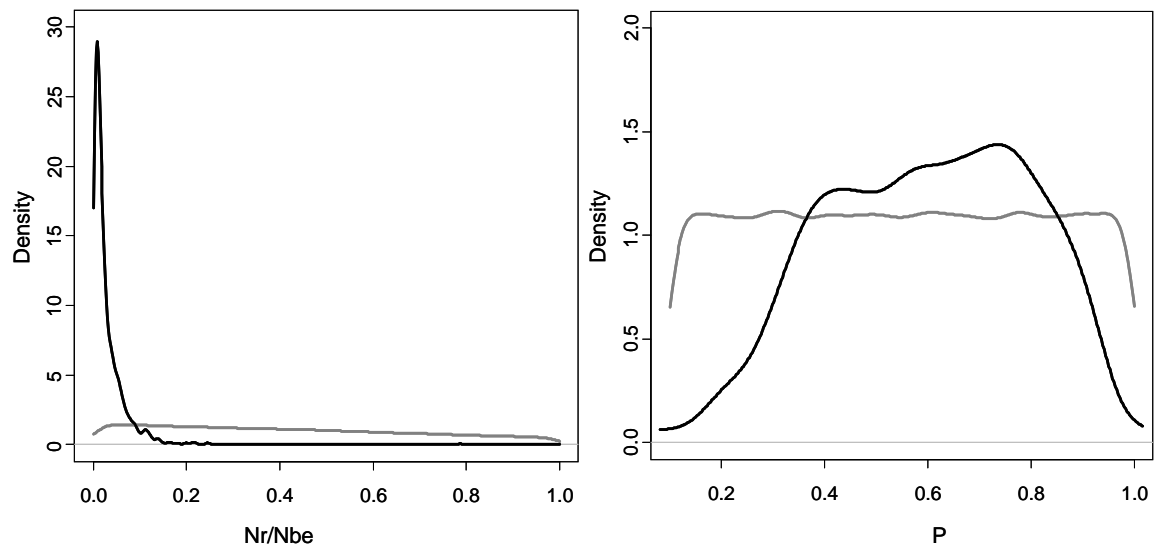
**Table S2.** Prior setting for demographic scenario of *E. urophylla* in Timor using the Approximate Bayesian Computation approach (Cornuet *et al.* 2008, 2010) and based on 17 SSR loci. SNM Standard Neutral Model (constant population size), BOT: bottleneck, EXP: demographic expansion, EXPBOT: recent expansion and past bottleneck and BOTEXP: recent bottleneck and past expansion. N, N1, N3, N4, Nb, Ne, Nbe, Nr, Nex, Neb define population sizes in the four scenario and tb, te, t1, t2, t3, t4 times of demographic event occurrence expressed in generation number.  $\mu$  mean is the average mutation rate and  $P$  mean is the average parameter of the Generalized Stepwise model geometric distribution. Locus specific mutation parameters ( $\mu_i$  and  $P_i$ ), considered as nuisance parameters, were kept at default values.

	Parameter	Distribution	Prior interval	
			Minimum	Maximum
SNM	N	Uniform	100	5.E+05
BOT	N1	Uniform	100	5.E+05
	Nb	Uniform	100	5.E+05
	tb	Uniform	1	1.E+04
EXP	N2	Uniform	100	5.E+05
	Ne	Uniform	100	5.E+05
	te	Uniform	1	1.E+04
EXPBOT	N3	Uniform	100	5.E+05
	Nr	Uniform	100	5.E+05
	Nbe	Uniform	100	5.E+05
	t1	Uniform	1	1.E+04
	t2	Uniform	1	1.E+04
BOTEXP	N4	Uniform	100	5.E+05
	Nex	Uniform	100	5.E+05
	Neb	Uniform	100	5.E+05
	t3	Uniform	1	1.E+04
	t4	Uniform	1	1.E+04
All scenario	$\mu_{\text{mean}}$	Uniform	1E-04	1E-03
	$P_{\text{mean}}$	Uniform	1E-01	5E+00



**Figure S1.** Density function for parameter prior (grey) and posterior (black) distributions for scenario 3.  $N_e$  and  $N_2$  define population sizes and  $T_e$  expansion time expressed in generation number.  $\mu_{micA}$  is the average mutation rate and  $P_{mean}$  is the average parameter of the Generalized Stepwise model geometric distribution. Note that the effective prior of  $N_2$  and  $N_e$  are not uniforms because of the constraint  $N_e > N_2$  we apply on these parameters. We also represented  $N_e/N_2$  which represent the inverse of population expansion.





**Figure S2.** Density function for parameter prior (grey) and posterior (black) distributions for scenario 4  $N3$ ,  $Nr$  and  $N3$  define population sizes.  $T1$  defines bottleneck time and  $T2$  defines expansion time expressed in generation number.  $M_{micA}$  is the average mutation rate and  $P_{mean}$  is the average parameter of the Generalized Stepwise model geometric distribution. Note that the effective prior of  $Nr$ ,  $Nbe$ ,  $N3$ ,  $T1$  and  $T2$  are not uniforms because of the constraints ( $Nbe > Nr$ ,  $Nr < N3$  and  $T1 < T2$ ) we apply on these parameters.



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